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<th>Description</th>
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</thead>
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<tr>
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<td>Operating instructions</td>
</tr>
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<td>General warning</td>
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<tr>
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<td>Caution</td>
</tr>
<tr>
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<td>Safe working load</td>
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<tr>
<td><img src="image" alt="Maximum patient weight" /></td>
<td>Maximum patient weight</td>
</tr>
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<td><img src="image" alt="In accordance with European Directive 2002/96/EC on Waste Electrical and Electronic Equipment, this symbol indicates that the product must not be disposed of as unsorted municipal waste, but should be collected separately. Refer to your local distributor for return and/or collection systems available in your country." /></td>
<td>In accordance with European Directive 2002/96/EC on Waste Electrical and Electronic Equipment, this symbol indicates that the product must not be disposed of as unsorted municipal waste, but should be collected separately. Refer to your local distributor for return and/or collection systems available in your country.</td>
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</tr>
<tr>
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<td>Date of manufacture</td>
</tr>
<tr>
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<td>No pushing</td>
</tr>
<tr>
<td><img src="image" alt="Do not store oxygen bottle" /></td>
<td>Do not store oxygen bottle</td>
</tr>
</tbody>
</table>
## Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>~</td>
<td>Alternating current</td>
</tr>
<tr>
<td></td>
<td>Direct current</td>
</tr>
<tr>
<td></td>
<td>Warning: non-ionizing radiation</td>
</tr>
<tr>
<td></td>
<td>Warning: crushing of hands</td>
</tr>
<tr>
<td></td>
<td>Dangerous voltage</td>
</tr>
<tr>
<td></td>
<td>Protective earth (ground)</td>
</tr>
<tr>
<td></td>
<td>Type B Applied Part</td>
</tr>
<tr>
<td>IPX6</td>
<td>Protection from powerful water jets</td>
</tr>
<tr>
<td>1 min / 20 min</td>
<td>Units of Duty Cycle - maximum time on is 1 minute / maximum time off is 20 minutes</td>
</tr>
</tbody>
</table>
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The words **WARNING**, **CAUTION**, and **NOTE** carry special meanings and should be carefully reviewed.

---

**WARNING**

Alerts the reader about a situation which, if not avoided, could result in death or serious injury. It may also describe potential serious adverse reactions and safety hazards.

---

**CAUTION**

Alerts the reader of a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the user or patient or damage to the equipment or other property. This includes special care necessary for the safe and effective use of the device and the care necessary to avoid damage to a device that may occur as a result of use or misuse.

---

**NOTE**

Provides special information to make maintenance easier or important instructions clearer.
This manual is designed to assist you with the operation of Stryker Model 1105 Prime Series® stretcher. Read this manual thoroughly before using the equipment or beginning maintenance on it. To ensure safe operation of this equipment, it is recommended that methods and procedures be established for educating and training staff on the safe operation of this stretcher.

PRODUCT DESCRIPTION

The Stryker Model 1105 Prime Series stretcher, with the retractable fifth wheel, optimizes traction and cornering to improve overall mobility.

INTENDED USE OF PRODUCT

The Stryker Model 1105 Prime Series stretcher is a wheeled device which consists of a platform mounted on a wheeled frame that is designed to support patients in a horizontal position. The device has siderails and has the option available to support the temporary or permanent placement of I.V. poles. A stretcher provides the operator with a method of transporting patients within a healthcare facility. The device has a fifth wheel that may be engaged to guide the stretcher along a straight line during transport and pivots the stretcher around corners. Some stretchers may also be used for minor procedures and short-term stay (treatment and recovery).

INTENDED USE OF PRODUCT - PRIME X OPTION

The Prime X option provides a platform for the short-term outpatient clinical evaluation and treatment of human patients and additionally may be used for minor procedures and short-term outpatient stay (treatment and recovery). The Prime X option provides an articulating radiographic patient support surface and a platform below the patient support surface for X-Ray cassette placement to allow the capture of clinical X-Rays (AP Full Body, optional Full Body Lateral, and optional Upright Chest) when used in conjunction with a medical X-Ray system. The Prime X option is a wheeled device consisting of a platform mounted on a wheeled frame that is designed to transport patients in a horizontal position within the interior of a healthcare facility by health professionals and/or trained representatives of the user facility. The device has siderails and optional supports for fluid infusion equipment.

The Prime X option is intended to be used to transport patients to and from all departments within the interior of a healthcare facility. The use of the Prime X option as a short-term outpatient clinical evaluation, treatment, minor procedure, and short-term outpatient recovery platform may include use in, but is not limited to, the Emergency Department (ED), including the Trauma area, and Post Anesthesia Care Unit (PACU). The Prime X option is not intended to be used for long-term inpatient treatment and recovery. See the specification table on page 10 for the intended environmental conditions.

The Prime X option has a safe working load up to 700 pounds (318 kg) and is intended to be used with all patients, including those mildly to critically ill. The stretcher may also be used to transport deceased patients within an enclosed healthcare facility.

The Prime X option has an optional scale system intended to measure and display weight in pounds or kilograms of patients weighing 50 to 700 pounds (22.7 to 318 kg) and patients not exceeding the height of 75.25 inches (191 cm). See the specification table for accuracy claims.

INTENDED USE OF PRODUCT - PRIME X OPTION (CONTINUED)

The Prime X option has the following options available: three-sided brake/steer pedal controls, three- and four-sided hydraulic lift controls, powered lift controls, dual siderail latch assembly, slider board assembly, pump bar, I.V. poles, restraint straps, defibrillator tray, defibrillator tray/foot extender, footboard/chart holder, serving tray, serving tray holder/footboard, siderail pads, upright oxygen bottle holder, and I.V. caddy and may include other options as described in this manual.
EXPECTED SERVICE LIFE

The Prime X option has an expected service life of 10 years under normal use, conditions, and with appropriate periodic maintenance as described in the maintenance manual for each device.

CONTRAINDICATIONS

The Prime X option is not recommended for use with a Stryker Pioneer mattress or a mattress with a thickness greater than four inches and is not compatible for use with a C-Arm.

The Prime X option is intended for use in all establishments other than home healthcare, domestic, and those directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.
## SPECIFICATIONS

<table>
<thead>
<tr>
<th></th>
<th>Prime 26” Width Option</th>
<th>Prime 30” Width Option</th>
<th>Prime X Option 30” Width</th>
</tr>
</thead>
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<tr>
<td><strong>Safe working load</strong></td>
<td>700 lb</td>
<td>700 lb</td>
<td>700 lb</td>
</tr>
<tr>
<td><strong>Note:</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>318 kg</td>
<td>318 kg</td>
<td>318 kg</td>
</tr>
<tr>
<td><strong>Maximum patient weight</strong></td>
<td>675 lb</td>
<td>675 lb</td>
<td>675 lb</td>
</tr>
<tr>
<td></td>
<td>306 kg</td>
<td>306 kg</td>
<td>306 kg</td>
</tr>
<tr>
<td><strong>Overall Stretcher Length</strong></td>
<td>85” (± .5”)</td>
<td>85” (± .5”)</td>
<td>85” (± .5”)</td>
</tr>
<tr>
<td></td>
<td>215.9 cm</td>
<td>215.9 cm</td>
<td>215.9 cm</td>
</tr>
<tr>
<td><strong>Overall Stretcher Width</strong></td>
<td>34” (± 1”)</td>
<td>38” (± 1”)</td>
<td>38” (± 1”)</td>
</tr>
<tr>
<td><strong>(Siderails Up)</strong></td>
<td>86.4 cm</td>
<td>96.5 cm</td>
<td>96.5 cm</td>
</tr>
<tr>
<td><strong>Overall Stretcher Width</strong></td>
<td>30.25” (± .5”)</td>
<td>30.5” (± .5”)</td>
<td>30.5” (± .5”)</td>
</tr>
<tr>
<td><strong>(Siderails Down)</strong></td>
<td>76.8 cm</td>
<td>77.5 cm</td>
<td>77.5 cm</td>
</tr>
<tr>
<td><strong>Minimum / Maximum</strong></td>
<td>20.75” / 34” (± 1”)</td>
<td>20.75” / 34” (± 1”)</td>
<td>23.25” / 36.5” (± 1”)</td>
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<tr>
<td><strong>Stretcher Height</strong></td>
<td>52.7 cm / 86.4 cm</td>
<td>52.7 cm / 86.4 cm</td>
<td>59.1 cm / 92.7 cm</td>
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<tr>
<td><strong>Fowler Angle</strong></td>
<td>0° to 90° (± 5°)</td>
<td></td>
<td></td>
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<tr>
<td><strong>Gatch Height</strong></td>
<td>5.5” (14 cm) minimum</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td><strong>Trendelenburg / Reverse</strong></td>
<td>+17°/-17° (± 3°)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Trendelenburg</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Minimum Under-Stretcher Clearance</strong></td>
<td>1.75” under the hydraulic jacks and fifth wheel</td>
<td>1.75” under the hydraulic jacks and fifth wheel</td>
<td>1.75” under the hydraulic jacks and fifth wheel</td>
</tr>
<tr>
<td></td>
<td>5.75” nominal</td>
<td>5.75” nominal</td>
<td>5.75” nominal</td>
</tr>
<tr>
<td></td>
<td>14.6 cm</td>
<td>14.6 cm</td>
<td>14.6 cm</td>
</tr>
<tr>
<td><strong>Attenuation Equivalent</strong></td>
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<td></td>
<td>Maximum value allowed is 1.7 mm Al</td>
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<tr>
<td><strong>(Aluminum Equivalence)</strong></td>
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### SPECIFICATIONS (CONTINUED)

<table>
<thead>
<tr>
<th>Electric Options</th>
<th>Optional Electric Litter (available on Prime Only)</th>
<th>Optional Electric Lift (available on Prime and Prime X Option)</th>
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<tr>
<td>Electrical Requirements</td>
<td>120V~, 60Hz, 10 A</td>
<td>120V~, 60Hz, 10 A</td>
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<tr>
<td>Duty Cycle</td>
<td>Continuous operation with intermittent loading is 1 min ON/20 min OFF</td>
<td>Continuous operation with intermittent loading is 1 min ON/20 min OFF</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional Scale System</th>
<th>Optional Electric Litter (available on Prime and Prime X Option)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery Type</td>
<td>Non-Electric Litter (available on Prime and Prime X Option)</td>
</tr>
<tr>
<td></td>
<td>Optional Electric Lift</td>
</tr>
<tr>
<td>Battery Type</td>
<td>4 x AA Battery (4 X 1.5V) Alkaline Type (LR6)</td>
</tr>
<tr>
<td></td>
<td>4 x AA Battery (4 X 1.5V) Alkaline Type (LR6)</td>
</tr>
<tr>
<td>Battery Voltage</td>
<td>6.0V~</td>
</tr>
<tr>
<td></td>
<td>6.0V~</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional Scale System</th>
<th>Optional Electric Litter</th>
<th>Optional Electric Lift</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery Type</td>
<td>1 x Rechargeable Lithium Ion Battery Pack (0058-135-000)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Battery Voltage</td>
<td>10.8V~, 2.4Ah</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional Scale System with Chaperone (Stretcher Exit)</th>
<th>Optional Electric Litter</th>
<th>Optional Electric Lift</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery Type</td>
<td>1 x Rechargeable Lithium Ion Battery Pack (0058-134-000)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Battery Voltage</td>
<td>10.8V~, 4.8 Ah</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

Stryker reserves the right to change specifications without notice.

**Note:** Equipment not suitable for use in the presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.

### STANDARDS APPLIED

<table>
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<tr>
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<th>Name</th>
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<tr>
<td>IEC 60601-1</td>
<td>Medical electrical equipment - Part 1: General requirements for basic safety and essential performance</td>
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<tr>
<td>IEC 60601-1-2</td>
<td>Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests</td>
</tr>
<tr>
<td>IEC 60601-1-3</td>
<td>Medical electrical equipment - Part 1-3: General requirements for basic safety and essential performance - Collateral Standard: Radiation protection in diagnostic X-Ray equipment</td>
</tr>
<tr>
<td>IEC 60601-2-52</td>
<td>Medical electrical equipment - Part 2-52: Particular requirements for the basic safety and essential performance of medical beds</td>
</tr>
<tr>
<td>IEC 60601-2-54</td>
<td>Medical electrical equipment - Part 2-54: Particular requirements for the basic safety and essential performance of X-Ray equipment for radiography and radioscopy</td>
</tr>
</tbody>
</table>
### SPECIFICATIONS (CONTINUED)

<table>
<thead>
<tr>
<th>Environmental Conditions</th>
<th>Operation</th>
<th>Storage and Transportation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Temperature</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100°F (38°C)</td>
<td>140°F (60°C)</td>
<td></td>
</tr>
<tr>
<td>50°F (10°C)</td>
<td>-4°F (-20°C)</td>
<td></td>
</tr>
<tr>
<td><strong>Relative Humidity</strong></td>
<td>75%</td>
<td>95%</td>
</tr>
<tr>
<td>30%</td>
<td>10%</td>
<td></td>
</tr>
<tr>
<td><strong>Atmospheric Pressure</strong></td>
<td>1060 hPa</td>
<td>1060 hPa</td>
</tr>
<tr>
<td>700 hPa</td>
<td>500 hPa</td>
<td></td>
</tr>
</tbody>
</table>

Specifications listed are approximate and may vary slightly from unit to unit or by power supply fluctuations.

### SPECIFICATIONS - OPTIONAL SCALE SYSTEM (NON-ELECTRIC LITTER/ELECTRIC LITTER OPTION)

- **Optional Scale System Weight Operating Range**: 50 lb (22.7 kg) to 700 lb (318 kg)
- **Optional Scale System Accuracy**:
  - ±3 lb (1.3 kg) for weights less than 100 lb (45 kg) and ±3% for weights greater than or equal to 100 lb (45 kg) *

<table>
<thead>
<tr>
<th>Environmental Conditions</th>
<th>Operation</th>
<th>Storage and Transportation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Temperature</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>79°F (26°C)</td>
<td>140°F (60°C)</td>
<td></td>
</tr>
<tr>
<td>61°F (16°C)</td>
<td>-4°F (-20°C)</td>
<td></td>
</tr>
<tr>
<td><strong>Relative Humidity</strong></td>
<td>75%</td>
<td>95%</td>
</tr>
<tr>
<td>30%</td>
<td>10%</td>
<td></td>
</tr>
<tr>
<td><strong>Atmospheric Pressure</strong></td>
<td>1060 hPa</td>
<td>1060 hPa</td>
</tr>
<tr>
<td>700 hPa</td>
<td>500 hPa</td>
<td></td>
</tr>
</tbody>
</table>

* To meet this accuracy claim, the patient surface must be in the flat position (fowler and gatch down) and the stretcher cannot exceed 5 degrees of Trendelenburg/reverse Trendelenburg.
Introduction

PRODUCT ILLUSTRATION - ELECTRIC LITTER OPTION

Figure 1: Electric Litter Option
Introduction

PRODUCT ILLUSTRATION - PRIME X OPTION

Figure 2: Prime X Option
Introduction

Figure 3: Type B Applied Parts - Prime
Introduction

PRODUCT ILLUSTRATION - TYPE B APPLIED PARTS - PRIME X OPTION

Figure 4: Type B Applied Parts - Prime X Option
CONTACT INFORMATION

Contact Stryker Customer Service or Technical Support at: (800) 327-0770 or (269) 324-6500.

Stryker Medical
3800 E. Centre Avenue
Portage, MI 49002
USA

Please have the serial number (A) of your Stryker product available when calling Stryker Customer Service or Technical Support. Include the serial number in all written communication.

SERIAL NUMBER LOCATION

Figure 5: Serial Number Location
Specification Labels

1105-101-002

1105-101-003

1105-101-006

1105-101-007

1105-101-008
Summary of Safety Precautions

Carefully read and strictly follow the warnings and cautions listed within this manual. Service only by qualified personnel. See the maintenance manual for additional information.

⚠️ WARNING

- This stretcher is equipped with a hospital grade plug for protection against electric shock hazard. It must be plugged directly into a properly grounded three-prong receptacle. Grounding reliability can be achieved only when a hospital grade receptacle is used.
- Always apply the brakes when a patient is getting on or off the stretcher. Push on the stretcher to ensure that the brakes are securely locked. Always engage the brakes unless the stretcher is being moved. Injury could result if the stretcher moves while a patient is getting on or off the stretcher.
- Patients should be discouraged from sitting directly on the ends of the stretcher. Excessive weight could cause the litter surface to tip up, possibly causing patient injury.
- Leave the stretcher height in the lowest position when the patient is left unattended. Leaving the stretcher height in a raised position could increase the chance of patient falls and injury.
- Make sure that the brakes are completely released before attempting to move the unit. Attempting to move the unit with the brakes engaged could result in injury to the user and/or patient.
- After raising the siderails, pull firmly on the siderail to ensure that it is securely locked in the up position. Siderails are not intended to serve as a patient restraint device to keep patients from exiting the unit. Siderails are intended to keep a patient from inadvertently rolling off the unit. It is the responsibility of the attending medical personnel to determine the degree of restraint necessary to ensure that a patient will remain in place. Failure to utilize the siderails properly could result in patient injury.
- When lowering the siderail to the collapsed position, keep extremities of patients and staff away from the siderail spindles or injury could occur.
- Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.
- Operation of the fowler is a manual procedure. Use caution when raising the fowler while a patient is on the stretcher. Use proper lifting techniques and get additional assistance, if necessary. Failure to use proper lifting techniques could cause injury to the operator.
- Keep hands/fingers clear of the area around the fowler release handle and the fowler frame when lowering the fowler. Injury could result if care is not taken when lowering the fowler.
- To avoid the risk of injury, ensure that the gatch prop rod is fully raised and securely placed in position.
- Use caution when operating the gatch while a patient is on the stretcher. Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.
- Use caution when operating the recovery chair while a patient is on the stretcher. Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.
- To avoid patient injury or equipment damage, all lines from any equipment stored on the pump rack must be diverted away from the gatch handles.
- To avoid patient injury or equipment damage, do not lift the stretcher by the pump rack.
- To avoid equipment damage, remove any equipment from the pump rack that may be in the way before lowering the litter.
- To avoid equipment damage while transporting the stretcher, verify that any equipment on the pump rack can safely pass through door openings and under light fixtures.
- When using the Prime X option in conjunction with devices that generate X-radiation, the generating devices may produce residual, stray, and/or scattered radiation. Users should refer to local, state, and federal use guidelines, as well as appropriate facility protocols, for safety before use. Special attention should be given when performing X-Rays with the stretcher’s fowler in the upright position and also when performing X-Rays using a lateral cassette.
- The Prime X option is not recommended for use with a Stryker Pioneer mattress or a mattress with a thickness greater than four inches and is not compatible for use with a C-Arm.
Summary of Safety Precautions

WARNING (CONTINUED)

• To avoid the risk of patient or operator injury, ensure that all devices placed on the defibrillator tray are securely strapped to the tray.
• To avoid the risk of patient or operator injury, ensure that all devices placed on the foot extender/defibrillator tray are securely strapped to the tray.
• To avoid the risk of injury to the patient or user or damage to the I.V. pole while transporting the stretcher, make sure that the I.V. caddy is securely tightened on the I.V. pole.
• To avoid the risk of patient injury or equipment damage, do not sit on the foot support.
• Physical restraints, even if properly secured, may result in serious harm to patients and caregivers. The use of restraint straps may potentially cause entanglement, entrapment, physical injury, and/or death. Caution must be used in affixing restraint straps to avoid potential injury to both patients and caregivers.
• Restraint straps and/or devices must be attached only at the identified attachment points of the unit. Failure to do so may result in patient or caregiver injury. Do not attach restraints straps to the siderail.
• This unit accommodates the use of ankle, chest, wrist, and body restraints. The use of restraint straps is regulated by state and federal restrictions. Users, caregivers, and/or practitioners should refer to the applicable state and federal restrictions and the appropriate facility protocols before using any restraint strap and/or device.
• If your unit is equipped with the optional electric lift/litter, unplug the power cord from the power outlet before transporting or cleaning the unit.
• This device does not offer any protection against X-Ray radiation.
• Do not steam clean the unit.
• Medical electrical equipment (such as the optional scale system or optional electric lift/litter) requires special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided on page 40 to prevent equipment malfunction.
• Portable and mobile RF communication equipment can affect Medical Electrical Equipment (such as the optional scale system or optional electric lift/litter).
• To avoid malfunction, the optional scale system or optional electric lift/litter should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the optional scale system or optional electric lift/litter should be observed to verify normal operation in the configuration in which it will be used.
• When using any mattress thicker than 2.5 inches or when using a mattress overlay with the Prime X option, extra caution and operator supervision is recommended to reduce the risk of patient falls due to lesser siderail coverage.
• Do not immerse mattress in cleaning or disinfectant solutions. Excess moisture could cause equipment malfunction resulting in equipment damage or patient injury.
• Do not allow fluid to pool on the mattress. Fluids can cause corrosion of components and may cause the safety and performance of this equipment to become unpredictable.
• Inspect mattress covers for tears, punctures, excessive wear, and misaligned zippers each time the covers are cleaned. If compromised, the mattress should be removed from service immediately and replaced to prevent cross-contamination.
• Disinfect the mattress between patients. Failure to do so could result in cross-contamination and infection.
• Some disinfectants may cause damage to the product if used improperly. If the products described below are used to disinfect the mattress, measures must be taken to ensure the entire surface is wiped with a damp cloth soaked in clean water and thoroughly dried following disinfection. The cover can be damaged when exposed to such disinfectants beyond the manufacturers’ recommendations. Failure to follow these directions when using these types of disinfectants may void this product warranty.
• The mattress cover must be completely dry before storage or adding linens. Failure to remove excess disinfectant could cause degradation of the cover material.
Summary of Safety Precautions

⚠️ CAUTION

- This stretcher is not intended for pediatric use or for patients under 50 lb. This stretcher is intended for use by trained hospital personnel only.
- Do not modify this stretcher. Modifying the unit can cause unpredictable operation resulting in injury to the patient or operator. Modifying the unit will also void its warranty.
- To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
- Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.
- To avoid injury or damage to the equipment, do not allow the siderail to lower on its own.
- The weight capacity of the gatch is 200 lb. Do not sit or stand on the gatch. Injury or damage to the equipment could occur.
- To achieve recovery chair position, your stretcher must be equipped with the Lift Assist™ backrest and gatch options.
- The weight capacity of the base hood is 60 lb. Do not sit or stand on the base hood. Injury or damage to the equipment could occur.
- Do not step on the base hood.
- Do not use the cutout for the oxygen bottle holder on the base hood for the storage of oxygen bottles or patient belongings.
- The weight capacity of the pump rack is 40 lb.
- Do not use the pump rack as a push/pull device because equipment damage could occur.
- To avoid damage, do not put items weighing more than 30 lb on the defibrillator tray.
- Do not use the defibrillator tray as a push/pull device because equipment damage could occur.
- If the stretcher is equipped with the optional foot end I.V. pole, the I.V. pole must be in the raised position when the foot extension/defibrillator tray is installed. If the I.V. pole is not raised, the foot extension will not function properly and injury could occur.
- The push handles were designed for use while transporting the stretcher. Avoid using other parts of the stretcher as push/pull devices unless otherwise specified in the manual because damage could occur.
- To avoid damage, do not put items weighing more than 30 lb on the foot extender/defibrillator tray.
- Do not use the foot extension/defibrillator tray as a push/pull device because equipment damage could occur.
- Do not use the footboard/chartholder as a push/pull device because equipment damage could occur.
- Always store the I.V. caddy when not in use to avoid damaging it when the unit is moved.
- To avoid the risk of equipment damage, do not use the foot support to store patient belongings or other items.
- To avoid injury to the operator, ensure that the operator’s fingers are clear of the mechanism when positioning the foot support.
- Foot supports should be in the stored position when moving. The stretcher should be in brake position when foot supports are in use.
- To avoid the risk of damage to the equipment, do not use the foot support as a push/pull device.
- To avoid injury to the patient or operator, ensure foot supports are tightened securely prior to use.
- If the stretcher is equipped with the scale system option, the scale should not be utilized while the foot supports are in use because inaccurate readings may occur.
- If the stretcher is equipped with the chaperone option, the chaperone option should not be utilized while the foot supports are in use because false readings may occur.
- To avoid damage to the removable I.V. pole, the weight of the I.V. bags should not exceed 40 lb.
- To avoid damage, the safe working load of the two-stage permanently attached I.V. pole is 40 lb.
- To avoid damage while transporting the stretcher, verify that the I.V. pole is at a low enough height to allow it to safely pass through door openings and under light fixtures.
Summary of Safety Precautions

⚠️ CAUTION (CONTINUED)

- To avoid damage, the weight of the I.V. bags should not exceed 12 lb while the weight of any one item attached to each stage of the three-stage permanently attached I.V. pole should not exceed 9.3 lb.
- Do not use the I.V. pole as a push/pull device because equipment damage could occur.
- To avoid damage, do not put items weighing more than 40 lb in the upright oxygen bottle holder.
- Do not use the upright oxygen bottle holder as a push/pull device because equipment damage could occur.
- To avoid damage, do not put items weighing more than 30 lb on the serving tray.
- To avoid risk of user injury or damage to the equipment, ensure that the Upright X-Ray Cassette Holder is installed correctly following the instructions in “Using the Optional Upright X-Ray Cassette Holder (Prime X Option Only)” on page 64.
- To avoid risk of user injury or damage to the equipment, ensure that the Lateral X-Ray Cassette Holder is installed correctly following the instructions “Using the Optional Lateral Cassette Holder (Prime X Option Only)” on page 65.
- Do not use the serving tray holder/footboard as a push/pull device because equipment damage could occur.
- Do not iron, dry-clean, or tumble dry the mattress, as this will cause malfunction and damage the product.
- The mattress cover must be completely dry before storing, adding linens or placing a patient on the surface, to prevent the performance of the equipment from being impaired.
- Avoid over exposure to alcohol or hydrogen peroxide. Swelling of the cover material will result.
- Do not allow liquid to seep into the zipper area and watershed cover barrier. Fluids allowed to come in contact with the zipper may leak into the mattress which could impair the equipment performance.
- Frequent or prolonged exposure to higher concentration disinfectant solutions may prematurely age the cover fabric.
- The use of accelerated hydrogen peroxides or quaternaries containing glycol ethers may damage the cover.
- Before returning the unit to service after cleaning, verify that labels are intact, raise/lower the stretcher, lock the brake/steer pedal in both positions, latch/unlatch the siderails, and raise/lower the Fowler and gatch.

NOTE

- Equipment not suitable for use in the presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.
- The bottom of the brake pads should be cleaned regularly to prevent wax or floor remnant buildup.
- Clean the base hood storage area regularly.
- In lieu of specific requirements from IEC 60601-1 and IEC 60601-2-54 equivalent means of safety were used on the Prime-X series of products regarding accessory overloading, applied parts, and indications to the operator.
- There is a dual siderail latch option available with latches on both ends of the stretcher.
Summary of Safety Precautions

PINCH POINTS - PRIME X OPTION

Figure 6: Pinch Points - Prime X Option
Setup Procedures

If this unit is equipped with the optional electric lift/litter, the unit must reach room temperature prior to conducting any setup and/or unit operations to prevent permanent damage to the unit.

Make sure that the unit is working properly before it is put into service. The following list will ensure that each part of the unit is checked.

Stretcher checklist:

1. Depress the pedal at either end of the stretcher fully to set the four wheel brakes and verify that all of the four casters are locked (page 25).
2. Raise and lower the hydraulic lift system (page 26 or page 27).
3. Raise the unit completely and activate the Trendelenburg function. Ensure that the head end lowers to the full down position (page 29).
4. Raise the unit completely and activate the reverse Trendelenburg function. Ensure that the foot end lowers to the full down position (page 29).
5. Run through the operation of the fifth wheel to ensure that it is operating properly (page 30).
6. Ensure that the siderails raise and lower smoothly and lock securely in the full up position (page 32).
7. Raise and lower the fowler (head end) (page 38).
8. Raise and lower the gatch (foot end) (page 40).

If equipped with the optional electric lift/litter:

1. Check all items on the stretcher checklist above.
2. Plug the unit into a properly grounded, hospital grade wall receptacle and ensure that the LED lights illuminate on the lockout keypad.
3. Raise and lower the hydraulic lift system (page 28).
4. Perform each function on the patient siderail controls to ensure that they are working properly (page 33).
5. Perform each function on the foot end nursing controls to ensure that they are working properly (page 34).
6. Raise and lower the fowler (head end) (page 39).
7. Raise and lower the gatch (foot end) (page 41).

If equipped with the optional scale - electric litter option (with or without chaperone option):

1. Check all items on the stretcher checklist above.
2. Plug the power cord into a properly grounded, hospital grade wall receptacle to charge the batteries.

Note: To charge the battery, see “Charging the Optional Scale System Battery Pack - Electric Litter Option (Not Available With Prime X Option)” on page 50 or “Charging the Optional Scale System Battery Pack - Electric Litter Option with Chaperone (Not Available With Prime X Option)” on page 52.

WARNING
This stretcher is equipped with a hospital grade plug for protection against electric shock hazard. It must be plugged directly into a properly grounded three-prong receptacle. Grounding reliability can be achieved only when a hospital grade receptacle is used.

CAUTION
• Do not modify this stretcher. Modifying the unit can cause unpredictable operation resulting in injury to the patient or operator. Modifying the unit will also void its warranty.
• In order to isolate the Prime Series stretcher from supply mains, disconnect electrical connection from the wall socket.
• Do not position the Prime Series stretcher in such a way as to restrict access to the wall plug.
• This stretcher is not intended for pediatric use or for patients under 50 lb. This stretcher is intended for use by trained hospital personnel only.
APPLYING THE BRAKE SYSTEM

For user convenience, a brake/steer control pedal is located on both ends of the stretcher as shown in Figure 7.

WARNING

Always apply the brakes when a patient is getting on or off the stretcher. Push on the stretcher to ensure that the brakes are securely locked. Always engage the brakes unless the stretcher is being moved. Injury could result if the stretcher moves while a patient is getting on or off the stretcher.

**To engage the brakes on the head end,** push down on the brake (red) side of pedal (A).
**To engage the brakes on the foot end,** push down on the brake (red) side of pedal (B).

**To release the brakes on the head end,** push down on the steer (green) side of pedal (A).
**To release the brakes on the foot end,** push down on the steer (green) side of pedal (B).

Note: Your stretcher may be equipped with optional side control brake and steer functions (C) in addition to the standard head end (A) and foot end (B) controls. The side control brakes operate the same as the head end and foot end brakes.

Note: The bottom of the brake pads should be cleaned regularly to prevent wax or floor remnant buildup.
OPERATING THE BASE CONTROLS - SIDE CONTROL HYDRAULICS

To operate the base controls, see Figure 9 to locate which pedals are used for what operation.

⚠️ CAUTION

- To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
- Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.

To raise the litter height, pump pedal (A) repeatedly until the desired height is achieved.

To lower both ends of the litter together, depress the center of pedal (B).

To lower only the head end of the litter, depress the side of pedal (B) closest to the head end of the stretcher.

To lower only the foot end of the litter, depress the side of pedal (B) closest to the foot end of the stretcher.

⚠️ WARNING

- Patients should be discouraged from sitting directly on the ends of the stretcher. Excessive weight could cause the litter surface to tip up, possibly causing patient injury.
- Leave the stretcher height in the lowest position when the patient is left unattended. Leaving the stretcher height in a raised position could increase the chance of patient falls and injury.
OPERATING THE BASE CONTROLS - OPTIONAL THREE-SIDED OR FOUR-SIDED CONTROL HYDRAULICS

To operate the base controls, see Figure 11 to locate which pedals are used for what operation.

⚠️ **CAUTION**

- To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
- Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.

**To raise the litter height**, pump pedal (A) repeatedly until the desired height is achieved.

**To lower both ends of the litter together**, depress pedal (B) and pedal (D) together using the same foot or depress in the center of pedal (C).

**To lower the foot end of the litter**, depress pedal (B) or the side of pedal (C) closest to the foot end.

**To lower the head end of the litter**, depress pedal (D) or the side of pedal (C) closest to the head end.

⚠️ **WARNING**

- Patients should be discouraged from sitting directly on the ends of the stretcher. Excessive weight could cause the litter surface to tip up, possibly causing patient injury.
- Leave the stretcher height in the lowest position when the patient is left unattended. Leaving the stretcher height in a raised position could increase the chance of patient falls and injury.
RAISING AND LOWERING THE LITTER HEIGHT - OPTIONAL ELECTRIC LIFT

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric lift.

⚠️ CAUTION

• To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
• Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.

To raise the litter height electrically, depress pedal (A) (see Figure 9 on page 26 or Figure 11 on page 27). The litter will begin to raise. Hold the pedal down until the desired litter height is achieved. Release the pedal at any time to stop the litter motion.

To lower the litter height manually, see the Operating the Base Controls - Side Control Hydraulics information on page 26 or Operating the Base Controls - Optional Three- or Four-Sided Hydraulic Controls information on page 27. The litter height does not lower electrically.

⚠️ WARNING

• Patients should be discouraged from sitting directly on the ends of the stretcher. Excessive weight could cause the litter surface to tip up, possibly causing patient injury.
• Leave the stretcher height in the lowest position when the patient is left unattended. Leaving the stretcher height in a raised position could increase the chance of patient falls and injury.
ADJUSTING TRENDELENBURG/REVERSE TRENDELENBURG POSITIONS - SIDE CONTROL HYDRAULICS

Litter height must first be raised in order to achieve a Trendelenburg or reverse Trendelenburg position.

⚠️ CAUTION
• To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
• Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.

For Trendelenburg positioning (head down), depress the side of pedal (B) closest to the head end (see Figure 9 on page 26).

For reverse Trendelenburg positioning (foot down), depress the side of pedal (B) closest to the foot end (see Figure 9 on page 26).

To lower the stretcher from reverse Trendelenburg position, depress pedal (A) once to raise the foot end of the stretcher, and then depress pedal (B) (see Figure 9 on page 26).

Note: The higher the litter is before pedal (B) is activated, the greater the Trendelenburg or reverse Trendelenburg angle will be. (Maximum Trendelenburg angle is +17°. Maximum reverse Trendelenburg angle is -17°.)

ADJUSTING TRENDELENBURG/REVERSE TRENDELENBURG POSITIONS - OPTIONAL THREE-OR FOUR-SIDED CONTROL HYDRAULICS

Litter height must first be raised in order to achieve a Trendelenburg or reverse Trendelenburg position.

⚠️ CAUTION
• To avoid damage, remove any equipment that may be in the way before raising or lowering the litter height.
• Do not raise the unit (hydraulics on base) with a patient lift under the stretcher.

For Trendelenburg positioning (head down) (see Figure 11 on page 27):
• Depress pedal (D) at the foot end of the unit or
• Depress the side of pedal (C) located on the patient left or patient right side, closest to the head end of the unit.

For reverse Trendelenburg positioning (foot down) (see Figure 11 on page 27):
• Depress pedal (B) at the foot end of the unit or
• Depress the side of pedal (C), located on the patient left or patient right side, closest to the foot end of the unit.

To lower the stretcher from reverse Trendelenburg position, depress pedal (A) once to raise the foot end of the stretcher and then depress pedal (C) or (D) (see Figure 11 on page 27).

Note: The higher the litter is before pedal (B), (C), or (D) is activated, the greater the Trendelenburg or reverse Trendelenburg angle will be. (Maximum Trendelenburg angle is +17°. Maximum reverse Trendelenburg angle is -17°.)
OPERATING THE FIFTH WHEEL

WARNING

• Make sure that the brakes are completely released before attempting to move the unit. Attempting to move the unit with the brakes engaged could result in injury to the user and/or patient.

• If your unit is equipped with the optional electric lift/litter, unplug the power cord from the power outlet before transporting or cleaning the unit. To unplug, grasp the mold near the outlet and pull the cord in a direction parallel to the floor (not at an angle).

The fifth wheel guides the stretcher along a straight line during transport and pivots the stretcher around corners.

To operate the fifth wheel, push the steer (green) side (A) of any brake/steer pedal to the lowest position as shown in Figure 14.

To disengage the fifth wheel, push the brake (red) side (B) of any brake/steer pedal to the neutral position as shown in Figure 14.
TRANSPORTING A PATIENT

**WARNING**

- Make sure that the brakes are completely released before attempting to move the product. Attempting to move the product with the brakes engaged could result in injury to the operator and/or patient.
- If your product is equipped with the optional electric lift/litter, unplug the power cord from the power outlet before transporting or cleaning the product.
- Do not transport the product laterally on inclines greater than 5.7 degrees (10% grade) to avoid tipping the product.

To transport a patient:

1. Put the litter in the lowest position. Make sure that the surface is flat.
2. Raise the siderails to the highest position.
3. If the product is equipped with the electric litter/electric lift, unplug the power cord from the wall outlet.
4. Push down on the steer (green) side of the brake/steer pedal.
5. Guide the product using the push handles.
OPERATING THE SIDERAIRS

Raising and lowering the siderails safely is a two-handed operation. Use one hand to hold and position the siderail and the other hand to operate the siderail latch.

To raise the siderails, pull up on the siderail (A) and raise it to the highest position until the latch (B) engages as shown in Figure 15.

⚠️ **WARNING**
- After raising the siderails, pull firmly on the siderail to ensure that it is securely locked in the up position. Siderails are not intended to serve as a patient restraint device to keep patients from exiting the unit. Siderails are intended to keep a patient from inadvertently rolling off the unit. It is the responsibility of the attending medical personnel to determine the degree of restraint necessary to ensure that a patient will remain in place. Failure to utilize the siderails properly could result in patient injury.
- Leave the stretcher height in the lowest position when the patient is left unattended. Leaving the stretcher height in a raised position could increase the chance of patient falls and injury.
- When using any mattress thicker than 2.5 inches or when using a mattress overlay with the Prime X option, extra caution and operator supervision is recommended to reduce the risk of patient falls due to lesser siderail coverage.

To lower the siderails, pull up on the latch (B) and guide the siderail to the lowest position as shown in Figure 15. The latches (B) are colored yellow for easy identification.

⚠️ **WARNING**
When lowering the siderail to the collapsed position, keep extremities of patients and staff away from the siderail spindles or injury could occur.

**Note:** The foot end of the siderail top rail can be used as a push/pull handle.

⚠️ **CAUTION**
To avoid injury or damage to the equipment, do not allow the siderail to lower on its own.

**Note:** There is a dual siderail latch option available with latches on both ends of the stretcher.
OPERATING THE SIDERAIL PATIENT CONTROLS - OPTIONAL ELECTRIC LITTER (NOT AVAILABLE WITH PRIME X OPTION)

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric litter.

Each siderail has backlit controls to allow the patient to position the fowler and gatch as shown in Figure 16. The power cord must be plugged into the wall socket for the patient controls to operate. When the stretcher is plugged in (powered) and the controls are unlocked (see page 35), the white buttons are illuminated.

Note: The siderail patient controls are positioned in a staggered location on each side of the stretcher for easy patient access.

**WARNING**

Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.

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**Figure 16: Siderail Patient Controls**

<table>
<thead>
<tr>
<th>Button</th>
<th>Button Name</th>
<th>Button Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Gatch Down</td>
<td>Press to lower the gatch (foot section)</td>
</tr>
<tr>
<td>2</td>
<td>Gatch Up</td>
<td>Press to raise the gatch (foot section)</td>
</tr>
<tr>
<td>3</td>
<td>Fowler Up</td>
<td>Press to raise the fowler (head section)</td>
</tr>
<tr>
<td>4</td>
<td>Fowler Down</td>
<td>Press to lower the fowler (head section)</td>
</tr>
</tbody>
</table>

---

- **Button Name**: Gatch Down
- **Button Function**: Press to lower the gatch (foot section)
- **Button Name**: Gatch Up
- **Button Function**: Press to raise the gatch (foot section)
- **Button Name**: Fowler Up
- **Button Function**: Press to raise the fowler (head section)
- **Button Name**: Fowler Down
- **Button Function**: Press to lower the fowler (head section)
OPERATING THE FOOT END NURSING CONTROLS - OPTIONAL ELECTRIC LITTER (NOT AVAILABLE WITH PRIME X OPTION)

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric litter.

The foot end nursing controls allow the operator to position the fowler and gatch as shown in Figure 17. The power cord must be plugged into the wall socket for the nursing controls to operate.

⚠️ WARNING
Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.

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<td>4</td>
<td>Fowler Down</td>
<td>Press to lower the fowler (head section)</td>
</tr>
</tbody>
</table>

Figure 17: Foot End Nursing Controls - Optional Electric Litter
USING PATIENT CONTROL LOCKOUT - OPTIONAL ELECTRIC LITTER (NOT AVAILABLE WITH PRIME X OPTION)

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric litter.

You can press the patient control lockout button to prevent the patient from using the siderail patient controls to move the fowler and gatch. The patient control lockout button is located at the foot end of the stretcher.

To lock the siderail patient controls, press the Lock/Unlock (A) button as shown in Figure 18. The lock icon (B) is illuminated amber while the patient controls are locked. The foot end nursing controls are not locked.

Note: When the siderail patient controls are locked, the siderail patient controls are not backlit.

To unlock the siderail patient controls, press the Lock/Unlock (A) button as shown in Figure 18. The unlock icon (C) is illuminated green when the patient controls are unlocked.
OPERATING THE OPTIONAL HEAD END PUSH HANDLES

To use the push handles, pivot the handles (A) up and push down until they are locked into position (Figure 19).
To store the push handles, lift the handles (B) up and pivot them down to store in the handle rests (Figure 20).

Figure 19: Head End Push Handles Open
Figure 20: Head End Push Handles Stored

⚠️ CAUTION

The push handles were designed for use while transporting the stretcher. Avoid using other parts of the stretcher as push/pull devices unless otherwise specified within this Operations Manual because equipment damage could occur.
OPERATING THE OPTIONAL FOOT END PUSH HANDLES

To use the push handles, pivot the handles (A) up and push down until they are locked into position (Figure 21).

To store the push handles, lift the handles (B) up and pivot them down to store in the handle rests (Figure 22).

![Figure 21: Foot End Push Handles Open](image1)

![Figure 22: Foot End Push Handles Stored](image2)

**CAUTION**
- The push handles were designed for use while transporting the stretcher. Avoid using other parts of the stretcher as push/pull devices unless otherwise specified in this Operations Manual because equipment damage could occur.
- To avoid the risk of patient and/or operator injury, raise the foot end push handles when using optional accessories (such as the foot extension/defibrillator tray, chart holder, upright oxygen bottle holder, or scale) or the accessories will not function properly.
- To avoid the risk of injury, keep fingers clear of the foot end push handles when lowering the optional gatch.
- If the stretcher is equipped with the optional foot end push handles, use caution if the foot extension/defibrillator tray, chart holder, and/or upright oxygen bottle holder is installed to avoid pinching your fingers.
OPERATING THE PNEUMATIC FOWLER - NON-ELECTRIC

To raise the fowler, squeeze either or both of the yellow fowler handles (A) for pneumatic assist until the fowler has reached the desired angle (between 0 and 90 degrees) as shown in Figures 23 and 24.

To lower the fowler, squeeze either or both of the yellow fowler handles (A) and push down until the fowler has reached the desired angle (between 90 and 0 degrees) as shown in Figures 23 and 24.

The drop seat/Lift Assist™ fowler uses the weight of the patient for additional assistance with raising the fowler. It also helps keep the patient from sliding toward the foot end of the stretcher when the fowler is raised.

WARNING

• Operation of the fowler is a manual procedure. Use caution when raising the fowler while a patient is on the stretcher. Use proper lifting techniques and get additional assistance, if necessary. Failure to use proper lifting techniques could cause injury to the operator.

• Keep hands/fingers clear of the area around the fowler release handle and the fowler frame when lowering the fowler. Injury could result if care is not taken when lowering the fowler.

Figure 23: Pneumatic Fowler

Figure 24: Pneumatic Fowler - Prime X Option
OPERATING THE FOWLER - OPTIONAL ELECTRIC LITTER (NOT AVAILABLE WITH PRIME X OPTION)

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric litter.

To raise the fowler, press the UP (3) button on the patient siderail controls (A) or foot end nursing controls (B) until the fowler has reached the desired angle (between 0 and 70 degrees) as shown in Figure 25.

To lower the fowler, press the DOWN (4) button until the fowler has reached the desired angle (between 70 and 0 degrees) as shown in Figure 25.

The drop seat/Lift Assist™ fowler uses the weight of the patient for additional assistance with raising the fowler. It also helps keep the patient from sliding toward the foot end of the stretcher when the fowler is raised.

**WARNING**

- Keep hands/fingers clear of the area around the fowler release handle and the fowler frame when lowering the fowler. Injury could result if care is not taken when lowering the fowler.
- Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.

![Figure 25: Fowler - Electric Option](image)
OPERATING THE OPTIONAL GATCH - NON-ELECTRIC (NOT AVAILABLE WITH PRIME X OPTION)

To raise the gatch, pump handle (B) repeatedly to the left until the gatch has reached the desired height (5.5”/14 cm minimum) as shown in Figure 28.

Note: You cannot raise the gatch manually if your unit is equipped with the optional electric litter.

To lower the gatch, pull handle (A) until the gatch has reached the desired height (5.5”/14 cm minimum) as shown in Figure 26.

CAUTION

The weight capacity of the gatch is 200 lb. Do not sit or stand on the gatch. Injury or damage to the equipment could occur.

To prop the foot end of the gatch up, lift up on the end of the gatch, allowing the prop rod to swing down and engage in the bracket as shown in Figure 28.

To release the prop, lift up on the end of the gatch, swing the prop rod toward the head end of the unit to disengage the bracket and lower the foot end as shown in Figure 28.

WARNING

To avoid the risk of injury, ensure that the gatch prop rod is fully raised and securely placed into position (Figure 28).
OPERATING THE GATCH - OPTIONAL ELECTRIC LITTER (NOT AVAILABLE WITH PRIME X OPTION)

Ensure that the power cord is plugged into a properly grounded, hospital grade wall outlet before using the optional electric litter.

**To raise the gatch**, press the UP (2) button on the siderail patient controls (A) or foot end nursing controls (B) until the gatch has reached the desired height (5.5”/14 cm minimum) as shown in Figure 29.

**To lower the gatch**, press the DOWN (1) button until the gatch has reached the desired height (5.5”/14 cm minimum) as shown in Figure 29.

---

**WARNING**

Use caution when operating the gatch while a patient is on the stretcher. Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.

---

**CAUTION**

The weight capacity of the gatch is 200 lb. Do not sit or stand on the gatch. Injury or damage to the equipment could occur.
OPERATING THE RECOVERY CHAIR (NOT AVAILABLE WITH PRIME X OPTION)

⚠️ CAUTION
To achieve recovery chair position, your stretcher must be equipped with the Lift Assist™ backrest and gatch options.

To place the stretcher into the recovery chair position as shown in Figure 30:
1. Raise the fowler to a seated position (for manual operation, see page 38; for the optional electric litter operation, see page 39).
2. Fully raise the gatch (for manual operation, see page 40; for the optional electric litter operation, see page 41).
3. Raise the stretcher to its highest height (for side control, see page 26; for three- or four-sided controls, see page 27; for optional electric litter operation, see page 28.)
4. Place the stretcher into the full reverse Trendelenburg position (see page 29).

![Figure 30: Recovery Chair](image)

To lower the stretcher from the recovery chair position:
1. Raise the stretcher to its highest height (for side control, see page 26 for three-or four-sided controls, see page 27; for optional electric litter operation, see page 28).
2. Lower the fowler from the seated position (for manual operation, see page 38; for the optional electric litter operation, see page 39).
3. Lower the gatch (for manual operation, see page 40; for the optional electric litter operation, see page 41).

⚠️ WARNING
Use caution when operating the recovery chair while a patient is on the stretcher. Powered stretcher mechanisms can cause serious injury. Operate stretcher only with persons clear of mechanisms.
USING THE BASE HOOD FOR STORAGE

You can store items in the base hood (A) as shown in Figure 31.

⚠️ CAUTION

- The weight capacity of the base hood is 60 lb. Do not sit or stand on the base hood. Injury or damage to the equipment could occur.
- Do not step on the base hood.
- Do not use the cutout for the oxygen bottle holder on the base hood for the storage of oxygen bottles or patient belongings.

Note: Clean the base hood storage area regularly.

Figure 31: Base Hood Storage
USING THE OPTIONAL PUMP RACK

**WARNING**

- To avoid patient injury or equipment damage, all lines from any equipment stored on the pump rack must be diverted away from the gatch handles.
- To avoid patient injury or equipment damage, do not lift the stretcher by the pump rack.
- To avoid equipment damage, remove any equipment from the pump rack that may be in the way before lowering the litter.
- To avoid equipment damage while transporting the stretcher, verify that any equipment on the pump rack can safely pass through door openings and under light fixtures.

**CAUTION**

- The weight capacity of the pump rack is 40 lb.
- Do not use the pump rack as a push/pull device because equipment damage could occur.

*Note:* The pump rack is an option that may have been installed at the foot end of the stretcher. The choice was made at the time that the stretcher was purchased.

The pump rack (A) can be used for the storage and transportation of stretcher equipment as shown in Figure 32.

![Figure 32: Pump Rack](image)
USING THE OPTIONAL RETRACTABLE CORD REEL - OPTIONAL ELECTRIC LIFT/LITTER

The retractable cord reel (A) stores the stretcher power cord during transport as shown in Figure 33.

To use the retractable cord reel:
1. Pull the cord out of the reel to the desired length.
2. Plug the power cord into a properly grounded, hospital grade wall outlet.

To store the power cord:
1. Unplug the plug by grasping the mold near the outlet and pull the cord in a direction parallel to the floor (not at an angle).
2. Tug and release the cord to retract the cord back into the cord reel.

WARNING
If your unit is equipped with the optional electric lift/litter, unplug the power cord from the power outlet before transporting or cleaning the unit. To unplug, grasp the mold near the outlet and pull the cord in a direction parallel to the floor (not at an angle).

Figure 33: Optional Retractable Cord Reel
OPERATING THE OPTIONAL SCALE SYSTEM

The scale option (see page 47) is available for units without the optional electric litter.

The scale option (see page 49) is available for units with the optional electric litter. The scale system has a battery backup option, so the standby icon indicates when the unit is unplugged and operating with battery backup.

The chaperone option (see page 53) is available for units with the optional electric litter scale option. The scale system with chaperone (stretcher exit) has a battery backup option, so the standby icon indicates when the unit is unplugged and operating with battery backup. The chaperone option also allows you to set zone controls to alert an operator when a patient may be attempting to exit the stretcher.
## OPERATING THE OPTIONAL SCALE SYSTEM - NON-ELECTRIC LITTER

![Diagram](image)

### Figure 34: Scale System Label - Non-Electric Litter

<table>
<thead>
<tr>
<th>Ref</th>
<th>Icon/Button</th>
<th>Description</th>
<th>Action</th>
<th>Display</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>Displays patient weight, unit of measurement, and battery status.</td>
<td></td>
<td>![Battery Icon]</td>
</tr>
<tr>
<td>2</td>
<td>![lb/kg]</td>
<td>Push to toggle between patient weight in pounds or weight in kilograms.</td>
<td>To convert the weight of the patient to kilograms, press and release <strong>lb/kg</strong>. Repeat to return to pounds.</td>
<td>XXX.X kg</td>
</tr>
<tr>
<td>3</td>
<td>![Weigh]</td>
<td>Push to weigh the patient. The display shows the patient’s weight for approximately 40 seconds before turning off.</td>
<td>Press and release <strong>Weigh</strong></td>
<td>XXX.X lb</td>
</tr>
<tr>
<td>4</td>
<td>![Zero]</td>
<td>Push and hold for 2 seconds to zero the scale system before putting a patient on the stretcher. If the display flashes “hold”, press and hold the <strong>Zero</strong> button again until the display reads “rel” (release). Release the <strong>Zero</strong> button. The display flashes “000.0”, then displays “000.0”. The system is not zeroed until the “000.0” stops flashing. For the most accurate results, always zero the scale system before putting a new patient on the stretcher. The display shuts off after approximately 40 seconds.</td>
<td>Press and hold <strong>Zero</strong> Release <strong>Zero</strong></td>
<td>hold rel 000.0 (flashing) 000.0 (solid)</td>
</tr>
</tbody>
</table>

**Note:** Do not touch the stretcher while the scale system is weighing or zeroing. The patient must remain still while the system is weighing. If the patient is moving, the system will try for 20 seconds to get a stable weight or zero value before displaying the error message **Err**.

If there is a loose connection or a malfunctioning component, the display will show “Err”. Attempt the function again. If the malfunction is still present, the display shows “Err” again. Call Stryker technical support at 800-327-0770.

To meet the accuracy claim as stated in the product specification on page 12, the patient surface must be in the flat position (fowler and gatch down) and the stretcher cannot exceed 5 degrees of Trendelenburg/reverse Trendelenburg.
REPLACING THE OPTIONAL SCALE SYSTEM BATTERIES - NON-ELECTRIC LITTER

To avoid completely draining the batteries and having the optional scale system shut down, replace the batteries whenever only one of the charge indicator bars on the display (1) is black as shown on page 47.

To replace the scale system batteries:
1. Remove the Phillips head screws that hold the battery compartment cover to the display assembly.
2. Replace all four AA batteries.
   • Install the positive and negative poles as indicated on the battery holder.
   • Use only Alkaline type (LR6) batteries.
   • Do not mix old and new batteries.
   • Properly dispose of the old batteries in accordance with local regulations.
3. Reinstall the screws and the cover.

If the display is flashing “Lo batt”, then the batteries are drained and the scale system is disabled. Replace the batteries with four new AA batteries as described above.
## Operating the Optional Scale System - Electric Litter Option Without Chaperone

(Not available with Prime X Option)

![Scale System Label - Electric Litter Option Without Chaperone](image)

### Figure 35: Scale System Label - Electric Litter Option Without Chaperone

<table>
<thead>
<tr>
<th>Ref</th>
<th>Icon/Button</th>
<th>Description</th>
<th>Action</th>
<th>Display</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>Displays patient weight, unit of measurement, and battery status.</td>
<td></td>
<td>![Battery Icon]</td>
</tr>
<tr>
<td>2</td>
<td>![lb/kg]</td>
<td>Push to toggle between patient weight in pounds or weight in kilograms.</td>
<td>To convert the weight of the patient to kilograms, press and release <code>lb/kg</code>. Repeat to return to pounds.</td>
<td>XXX.X kg XXX.X lb</td>
</tr>
<tr>
<td>3</td>
<td>![Zero]</td>
<td>Push and hold for 2 seconds to zero the scale system before putting a patient on the stretcher. If the display flashes “hold”, press and hold the <strong>Zero</strong> button again until the display reads “rel” (release). Release the <strong>Zero</strong> button. The display flashes “000.0”, then displays “000.0”. The system is not zeroed until the “000.0” stops flashing. For the most accurate results, always zero the scale system before putting a new patient on the stretcher. The display shuts off after approximately 40 seconds.</td>
<td>Press and hold <strong>Zero</strong> Release <strong>Zero</strong> hold rel 000.0 (flashing) 000.0 (solid)</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>![Power]</td>
<td>When the scale system is unplugged and operating with battery backup, the standby indicator is amber. When the unit is plugged in, the standby indicator is green.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>![Weigh]</td>
<td>Push to weigh the patient. The display shows the patient’s weight for approximately 40 seconds before turning off.</td>
<td>Press and release <strong>Weigh</strong></td>
<td>XXX.X lb</td>
</tr>
</tbody>
</table>

**Note:** Do not touch the stretcher while the scale system is weighing or zeroing. The patient must remain still while the system is weighing. If the patient is moving, the system will try for 20 seconds to get a stable weight or zero value before displaying the error message ![Error Icon](image). If there is a loose connection or a malfunctioning component, the display will show “Err”. Attempt the function again. If the malfunction is still present, the display shows “Err” again. Call Stryker technical support at 800-327-0770.

To meet the accuracy claim as stated in the product specification on page 12, the patient surface must be in the flat position (fowler and gatch down) and the stretcher cannot exceed 5 degrees of Trendelenburg/reverse Trendelenburg.
CHARGING THE OPTIONAL SCALE SYSTEM BATTERY PACK - ELECTRIC LITTER OPTION (NOT AVAILABLE WITH PRIME X OPTION)

To avoid completely draining the battery pack and having the optional scale system shut down, charge the battery pack whenever only one of the charge indicator bars on the display (1) is black as shown on page 49.

The battery pack charges whenever the power cord is plugged into a properly grounded, hospital grade power source. When the unit is stationary, plug the power cord into a power source whenever possible.

The optional scale system - electric litter option requires one 10.8V Li-Ion battery pack (0058-135-000). When fully discharged, the battery pack requires approximately 3 hours of charging time to recharge.
OPERATING THE OPTIONAL SCALE SYSTEM - ELECTRIC LITTER OPTION WITH CHAPERONE (NOT AVAILABLE WITH PRIME X OPTION)

Figure 36: Scale System Label - Electric Litter Option With Chaperone

<table>
<thead>
<tr>
<th>Ref</th>
<th>Icon/Button</th>
<th>Description</th>
<th>Action</th>
<th>Display</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>Indicates when Zone 1 is armed. Allows the patient to move around the stretcher freely, but cannot begin to exit the stretcher or the alert will sound.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>Displays patient weight, unit of measurement, and battery status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td>Push to toggle between patient weight in pounds or weight in kilograms</td>
<td>To convert the weight of the patient to kilograms, press and release lb/kg. Repeat to return to pounds.</td>
<td>XXX.X kg</td>
</tr>
<tr>
<td>4</td>
<td></td>
<td>Indicates when Zone 2 is armed. Zone 2 is more restrictive than Zone 1. When this zone is selected, the stretcher measures the patient’s center of gravity. If the patient’s center of gravity moves outside the preset boundary, an alert will sound.</td>
<td></td>
<td>XXX.X lb</td>
</tr>
</tbody>
</table>
| 5   |             | Push once to arm Zone 1. Push twice to arm Zone 2. Once armed or when alerting, press once to disarm. | Press and release Arm/Disarm | On 1  
Off |
| 6   |             | When the scale system is unplugged and operating with battery backup, the standby indicator is amber. When the unit is plugged in, the standby indicator is green. |        |         |
| 7   |             | Push to weigh the patient. The display shows the patient’s weight for approximately 40 seconds before turning off. | Press and release Weigh | XXX.X lb |
| 8   |             | Push and hold for 2 seconds to zero the scale system before putting a patient on the stretcher. If the display flashes “hold”, press and hold the Zero button again until the display reads “rel” (release). Release the Zero button. The display displays “000.0”, then displays “000.0”. The system is not zeroed until the “000.0” stops flashing. For the most accurate results, always zero the scale system before putting a new patient on the stretcher. The display shuts off after approximately 40 seconds. | Press and hold Zero  
Release Zero | hold rel  
000.0  
(ﬂashing)  
000.0  (solid) |
OPERATING THE OPTIONAL SCALE SYSTEM - ELECTRIC LITTER OPTION WITH CHAPERONE (NOT AVAILABLE WITH PRIME X OPTION) (CONTINUED)

**Note:** Do not touch the stretcher while the scale system is weighing or zeroing. The patient must remain still while the system is weighing. If the patient is moving, the system will try for 20 seconds to get a stable weight or zero value before displaying the error message "Err".

If there is a loose connection or a malfunctioning component, the display will show “Err”. Attempt the function again. If the malfunction is still present, the display shows “Err” again. Call Stryker technical support at 800-327-0770.

To meet the accuracy claim as stated in the product specification on page 12, the patient surface must be in the flat position (fowler and gatch down) and the stretcher cannot exceed 5 degrees of Trendelenburg/reverse Trendelenburg.

OPERATING THE CHAPERONE (STRETCHER EXIT) OPTION (NOT AVAILABLE WITH PRIME X OPTION)

To use the chaperone with zone control option:

1. Press the **Zero** button to reset the scale system.
   **Note:** Before positioning the patient on the stretcher, the scale system must be zeroed for the chaperone function to operate properly.
2. Position the patient on the stretcher and press the **Arm/Disarm** button to activate the chaperone function. The “Zone 1” LED will turn on. The chaperone function with zone control automatically selects Zone 1.
3. To select Zone 2 instead, press the **Arm/Disarm** button twice within three seconds of each other. The “Zone 2” LED will turn on.

To deactivate the chaperone function, press the **Arm/Disarm** button. The selected Zone light will turn off.

CHARGING THE OPTIONAL SCALE SYSTEM BATTERY PACK - ELECTRIC LITTER OPTION WITH CHAPERONE (NOT AVAILABLE WITH PRIME X OPTION)

To avoid completely draining the battery pack and having the optional scale system shut down, charge the battery pack whenever only one of the charge indicator bars on the display (2) is black as shown on page 51.

The battery pack charges whenever the power cord is plugged into a properly grounded, hospital grade power source. When the unit is stationary, plug the power cord into a power source whenever possible.

The optional scale system - with chaperone option only requires one 10.8V Li-Ion battery pack (0058-134-000). When fully discharged, the battery pack requires approximately 3 hours of charging time to recharge.
OPERATING THE CHAPERONE (STRETCHER EXIT) OPTION - OPTIONAL SETUP (NOT AVAILABLE WITH PRIME X OPTION)

To change the alert pattern:
1. Press and hold the Arm/Disarm (1) button and the Weigh (2) button together for 6 seconds. Ignore all display messages until “Ptrn” appears on the display (5).
2. Release both buttons. The display (5) shows the current setting “P (1-10)”.
3. Press the Arm/Disarm (1) or Weigh (2) button to change the setting. As you press each button to select your setting, a brief sample is played.
4. Press and hold the Arm/Disarm (1) button and the Weigh (2) button together for 6 seconds until “Set” appears on the display (5) to save your selected setting.
5. Release both buttons. The display (5) shows “P (1-10)”. A brief sample of your selected pattern will confirm your sound setting.

To change the alert volume:
1. Press and hold the Zero (3) button and the lb/kg (4) button together for 6 seconds. Ignore all display messages until “UOL” appears on the display (5).
2. Release both buttons. The display (5) shows the current setting “L (1-4)”.
3. Press the Zero (3) button or lb/kg (4) button to change the setting. As you press each button to select your setting, a brief sample is played.
4. Press and hold the Zero (3) button and the lb/kg (4) button together for 6 seconds until “Set” appears on the display (5) to save your selected setting.
5. Release both buttons. The display (5) shows “L (1-4)”. A brief sample of your selected volume will confirm your sound setting.
USING X-RAY CASSETTES - PRIME X OPTION

The Prime X option provides an articulating radiographic patient support surface and a platform below the patient support surface for X-Ray cassette placement to allow the capture of clinical X-Rays (AP Full Body, optional Full Body Lateral, and optional Upright Chest) when used in conjunction with medical X-Ray systems.

**WARNING**

When using the Prime X option in conjunction with devices that generate X-radiation, the generating devices may produce residual, stray, and/or scattered radiation. Users should refer to local, state, and federal use guidelines as well as appropriate facility protocols for safety before use. Special attention should be given when performing X-Rays with the stretcher’s fowler in the upright position and also when performing X-Rays using a lateral cassette.

You can insert and remove cassettes from the head end, foot end, and both sides of the stretcher as shown in Figure 38.

Center the patient on the stretcher by using the position indicator labels that are located on all sides of the stretcher.

**WARNING**

The Prime X option is not recommended for use with a Stryker Pioneer mattress or a mattress with a thickness greater than four inches and is not compatible for use with a C-Arm.

![Figure 38: Cassette Locations](image-url)
USING THE OPTIONAL DEFIBRILLATOR TRAY

To install the defibrillator tray, insert the pins on the defibrillator tray into the footboard sockets at the foot end of the stretcher as shown in Figure 39. Use the strap to secure the equipment to the tray.

**WARNING**

- To avoid risk of patient or operator injury, ensure that all devices placed on the defibrillator tray are securely strapped to the tray.
- Do not use the defibrillator tray as a push/pull device because equipment damage could occur.

**CAUTION**

To avoid damage, do not put items weighing more than 30 lb on the defibrillator tray.

USING THE OPTIONAL FOOT EXTENSION/DEFIBRILLATOR TRAY

To use as a defibrillator tray, pull out the top knob (A) and pivot the tray (B) over the foot extension (C) until the tray extends flat over the foot end of the stretcher as shown in Figure 38. Use the strap to secure the equipment to the tray.

**WARNING**

- To avoid risk of patient or operator injury, ensure that all devices placed on the defibrillator tray are securely strapped to the tray.
- Do not use the foot extension/defibrillator tray as a push/pull device because equipment damage could occur.

To use as a foot extension, pull out knob (A) and pivot the defibrillator tray back until it locks against the foot extension (C). While holding onto the assembly, pull out the bottom knob (D) and lower the foot extension down until it is flat as shown in Figure 40.

**CAUTION**

- If the stretcher is equipped with the optional foot end I.V. pole, the I.V. pole must be in the raised position when the foot extension/defibrillator tray is installed. If the I.V. pole is not raised, the foot extension will not function properly and injury could occur.
- If the stretcher is equipped with the optional foot end push handles, use caution if the foot extension/defibrillator tray, chart holder, and/or upright oxygen bottle holder is installed to avoid pinching your fingers.
- To avoid damage, do not put items weighing more than 30 lb on the foot extender/defibrillator tray.
USING THE OPTIONAL FOOTBOARD/CHARTHOLDER

To use the footboard/chart holder, insert the footboard/chart holder supports (A) into the corresponding holes located at the foot end of the stretcher.

⚠️ CAUTION

- Do not use the footboard/chart holder as a push/pull device because equipment damage could occur.
- If the stretcher is equipped with the optional foot end push handles, use caution if the foot extension/defibrillator tray, chart holder, and/or upright oxygen bottle holder is installed to avoid pinching your fingers.

USING THE OPTIONAL I.V. CADDY

To use the I.V. caddy:
1. Lift the I.V. caddy out of the storage tray or from the storage clip. Pivot the I.V. caddy to the desired position.
2. Turn knob (A) counterclockwise to loosen the pole clamp (C).
3. Pivot the knob (A) away from the clamp (B). The clamp (C) may then be opened.
4. Place the I.V. pole into the clamp (B). Close the clamp (C) around the I.V. pole and pivot the knob (A) back into position.
5. Turn the knob (A) clockwise to tighten it. The I.V. pole is ready to be transported with the unit.

To remove the I.V. pole from the I.V. caddy:
1. Turn knob (A) counterclockwise to loosen the pole clamp.
2. Pivot the knob away from the clamp (B), open the clamp and remove the I.V. pole from the I.V. caddy.

⚠️ CAUTION

Always store the I.V. caddy when not in use to avoid damaging it when the unit is moved.

⚠️ WARNING

To avoid the risk of injury to the patient or user or damage to the I.V. pole while transporting the stretcher, make sure that the I.V. caddy is securely tightened on the I.V. pole.
OPERATING THE OPTIONAL FOOT SUPPORTS (NOT AVAILABLE WITH PRIME X OPTION)

**WARNING**
To avoid the risk of patient injury or equipment damage, do not sit on the foot support.

To use the foot supports (see Figure 43):
1. Loosen the knee knob (A) at the top of the foot support to adjust the side-to-side angle of the foot support.
2. Secure the knee knob (A) to lock the foot support in the desired position.
3. Loosen the leg knob (B) on the side of the foot support to adjust the length.
4. Secure the leg knob (B) to lock the foot support in the desired position.
5. Flip the foot support (C) up before positioning patient.

The following options can be purchased, but cannot be utilized while the foot supports are in use:
- Pneumatic Backrest/Stationary Foot Composite
- Lift Assist™ Backrest/Stationary Foot Composite
- Pneumatic Backrest/Hydraulic Knee Gatch

To store the foot supports (see Figure 44):
1. Fully retract the foot supports using leg knob (B).
2. Fold the foot support pads (C) down toward the support bar.
3. Loosen the knee knob (A), rotate the foot supports to the position shown in the Figure 44 below.
4. Tighten to secure in place using the knee knob (A).
OPERATING THE OPTIONAL FOOT SUPPORTS (NOT AVAILABLE WITH PRIME X OPTION) (CONTINUED)

⚠️ CAUTION

- Do not use the foot support to store patient belongings or other items; equipment damage may occur.
- To avoid injury to the operator, ensure that the operator’s fingers are clear of the mechanism when positioning the foot support.
- Foot supports should be in the stored position when moving. The stretcher should be in brake position when foot supports are in use.
- To avoid the risk of damage to the equipment, do not use the foot support as a push/pull device.
- To avoid injury to the patient or operator, ensure foot supports are tightened securely prior to use.
- If the stretcher is equipped with the scale system option, the scale should not be utilized while the foot supports are in use; inaccurate readings may occur.
- If the stretcher is equipped with the chaperone option, the chaperone option should not be utilized while the foot supports are in use; false readings may occur.

The following options cannot be purchased if the foot support option is selected:
- Foot End Pop-Up Steering Handles
- Defibrillator Tray
- Defibrillator Tray/Foot Extender/Chart Service
- Serving Tray Holder/Footboard
- Footboard/Chart Holder
- Foot End I.V. Poles
OPERATING THE OPTIONAL TWO-STAGE PERMANENTLY ATTACHED I.V. POLE

**Note:** The two-stage permanently attached I.V. pole may have been installed at either the head, foot, or both ends of the stretcher. The choice was made at the time that the stretcher was purchased.

To use the two-stage permanently attached I.V. pole:
1. Lift and pivot the pole from the storage position and push down until it is locked into the receptacle.
2. To raise the height of the pole, pull up on the telescoping portion (A) until it locks into place at its fully raised position.
3. Rotate the I.V. hangers (B) to desired position and hang the I.V. bags.
4. To lower the I.V. pole, turn the latch (C) until section (A) lowers.

**CAUTION**
- To avoid damage, the safe working load of the two-stage permanently attached I.V. pole is 40 lb.
- To avoid damage while transporting the stretcher, verify that the I.V. pole is at a low enough height to allow it to safely pass through door openings and under light fixtures.

**Note:** The I.V. pole may be used as a push/pull device.
OPERATING THE OPTIONAL THREE-STAGE PERMANENTLY ATTACHED I.V. POLE

Note: The three-stage permanently attached I.V. pole may have been installed at either the head, foot, or both ends of the stretcher. The choice was made at the time that the stretcher was purchased.

To use the three-stage permanently attached I.V. pole:
1. Lift and pivot the pole from the storage position and push down until it is locked into the receptacle.
2. To raise the height of the pole, pull up on the telescoping portion (A) until it locks into place at its fully raised position.
3. For a higher I.V. pole, pull up on section (B). Release section (B) at any desired height and it will lock into place.
4. Rotate the I.V. hangers (C) to the desired position and hang the I.V. bags.
5. To lower the I.V. pole, push up on the red portion of the grip (D) while holding onto section (B) until it lowers. Turn the latch (E) until section (A) lowers.

CAUTION

- To avoid damage to the three-stage I.V. pole, the weight of the I.V. bags should not exceed 12 lb while the weight of any one item attached to each stage of the three-stage permanently attached I.V. pole should not exceed 9.3 lb.
- To avoid damage while transporting the stretcher, verify that the I.V. pole is at a low enough height to allow it to safely pass through door openings and under light fixtures.

Note: The I.V. pole may be used as a push/pull device.
OPERATING THE OPTIONAL REMOVABLE I.V. POLE

To use the removable I.V. pole:
1. Remove the I.V. pole from the storage trough under the litter and insert it into the receptacle on the corner of the litter frame.
2. To raise the height of the pole, turn the knob (B) counterclockwise and pull up on the telescoping portion (A) of the pole to raise it to the desired height.
3. Turn the knob (B) clockwise to lock the telescoping portion in place.

⚠️ CAUTION
- To avoid damage to the removable I.V. pole, the weight of the I.V. bags should not exceed 40 lb.
- To avoid damage while transporting the stretcher, verify that the I.V. pole is at a low enough height to allow it to safely pass through door openings and under light fixtures.
- Do not use the removable I.V. pole as a push/pull device because equipment damage could occur.

INSTALLING THE SIDERRAIL PADS

To install and use the siderail pads, tuck the siderail pad between the mattress and siderail. Then, attach the Velcro® straps around the top of the siderail to secure the pad to the siderail.

Figure 50: Removable I.V. Pole
USING THE OPTIONAL UPRIGHT OXYGEN BOTTLE HOLDER

To install the upright oxygen bottle holder, insert the support bar (A) into the I.V. socket at any of the four litter corners. Insert the cotter pin (B) through the hole in the support bar to hold the bottle holder in place as shown in Figure 51.

⚠️ CAUTION

- To avoid damage, do not put items weighing more than 40 lb in the upright oxygen bottle holder.
- Do not use the upright oxygen bottle holder as a push/pull device because equipment damage could occur.
- If the stretcher is equipped with the optional foot end push handles, use caution if the foot extension/defibrillator tray, chart holder, and/or upright oxygen bottle holder is installed to avoid pinching your fingers.

USING THE OPTIONAL SERVING TRAY

To use the serving tray, pull out on either end of the serving tray to extend it to the proper width to fit on top of the stretcher siderails as shown in Figure 52.

To store the serving tray in the optional serving tray holder/footboard, push in both ends of the serving tray and slide it into holder as shown in Figure 53.

⚠️ CAUTION

- To avoid damage, do not put items weighing more than 30 lb on the serving tray.
- Do not use the serving tray holder/footboard as a push/pull device because equipment damage could occur.
USING THE RESTRAINT STRAPS

This unit allows the use of ankle, chest, wrist, and body restraints. See Figures 54 and 55 for restraint strap attachment points. Do not attach restraints straps to the siderail. Stryker makes no recommendation for the use of restraints.

**WARNING**

- Physical restraints, even if properly secured, may result in serious harm to patients and caregivers. The use of restraint straps may potentially cause entanglement, entrapment, physical injury, and/or death. Caution must be used in affixing restraint straps to avoid potential injury to both patients and caregivers.
- Restraint straps and/or devices must be attached only at the identified attachment points of the unit. Failure to do so may result in patient or caregiver injury. Do not attach restraints straps to the siderail.
- This unit accommodates the use of ankle, chest, wrist, and body restraints. The use of restraint straps is regulated by state and federal restrictions. Users, caregivers, and/or practitioners should refer to the applicable state and federal restrictions and the appropriate facility protocols before using any restraint strap and/or device.

*Note:* The restraint straps are Type B applied parts.

Figure 54: Restraint Strap Location - Prime

Figure 55: Restraint Strap Location - Prime X
USING THE OPTIONAL UPRIGHT X-RAY CASSETTE HOLDER (PRIME X OPTION ONLY)

This unit stores the cassette against the backrest when raised in the upright position to help deliver superior image quality for chest X-Rays.

⚠️ WARNING

This device does not offer any protection against X-Ray radiation.

⚠️ CAUTION

To avoid risk of user injury or damage to the equipment, ensure that the Upright X-Ray Cassette Holder is installed correctly, following the instructions below.

To use the Upright X-Ray Cassette Holder

1. Raise the fowler section of the stretcher.
2. Position the upright cassette holder, placing the lower retainer guides (A) under the fowler weldment bar (Figure 56).
3. Raise the upright cassette holder until the hooks latch under the fowler weldment frame. Ensure that the upright cassette holder is securely latched.
4. Insert and remove cassettes from the upright cassette holder from either side of the cassette holder or pull up on the yellow arrow to release the wire latch (B) on the upright cassette holder, releasing the upright cassette holder from the fowler weldment, and then position the cassette in the holder.
5. Loosen the knob (E) on the back side of the upright cassette holder and adjust the cassette support rail (D) to position the cassette in the desired location (Figure 57). Tighten the knob to secure the cassette support rail in place.

To remove the Upright X-Ray Cassette Holder

1. With the fowler raised, pull up on the yellow arrow (B) to release the wire latch (C) on the upright cassette holder from the fowler weldment bar (Figure 56).
2. Lift up on the upright cassette holder to release the lower retainer guides (A) on the upright cassette holder from the fowler weldment bar (Figure 57).
**USING THE OPTIONAL LATERAL CASSETTE HOLDER (PRIME X OPTION ONLY)**

This removable cassette holder allows radiologists to take full length C-Spine images, holding virtually any size cassette.

**WARNING**

This device does not offer any protection against X-Ray radiation.

**CAUTION**

To avoid risk of user injury or damage to the equipment, ensure that the Lateral X-Ray Cassette Holder is installed correctly, following the instructions below.

**To use the Lateral Cassette Holder**

1. Press down on the yellow release button (A) to open the lateral cassette holder (Figure 58).
2. Slide the flat base (B) between the mattress and the patient platform (Figures 58 and 59).
3. Position the lateral cassette holder in the desired location.
4. Place the X-Ray cassette in the cassette canal (C) (Figure 59).
5. Pull up on the cassette hook (D) to adjust the height of the arm to fit snugly over the X-Ray cassette to hold the cassette in place (Figure 59).
6. Pull up on the cassette hook (D) to remove the X-Ray cassette from the lateral cassette holder (Figure 59).
7. Remove the lateral cassette holder from between the mattress and patient platform.
8. Press down on the release button to close the lateral cassette holder.
9. Store the lateral cassette holder as appropriate.

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**Figure 58: Lateral Cassette Holder**

**Figure 59: Lateral Cassette Holder**
CLEANING THE STRETCHER

**WARNING**

- If your unit is equipped with the optional electric lift/litter, unplug the power cord from the power outlet before transporting or cleaning the unit.
- Do not steam clean the unit.

These instructions are intended to provide recommended cleaning methods for Model 1105 Prime Series stretchers.

These units are designed to be power-washable. The unit may show some signs of oxidation or discoloration from continuous washing. However, no degradation of the stretcher’s performance characteristics or functionality will occur due to power washing as long as the proper procedures are followed.

**RECOMMENDED CLEANING METHOD**

- Follow the cleaning solution manufacturer’s dilution recommendations exactly.
- Stryker Medical recommends the standard hospital surgical cart washer for power washing.
- Do not replace the mattress on the stretcher until the unit is completely dry.

**CAUTION**

Before returning the unit to service after cleaning, verify that labels are intact, raise/lower the stretcher, lock the brake/steer pedal in both positions, latch/unlatch the siderails, and raise/lower the fowler and gatch.

**RECOMMENDED CART WASHING CLEANING METHOD**

Stryker Medical recommends using a standard hospital surgical cart washer to power wash the stretcher a maximum of once per year for the life of the unit.

To clean the unit with a cart washer:

1. Remove the mattress prior to washing the unit; do not wash the mattress with the stretcher.
2. Position the fowler at 45 degrees, position the gatch to the full down position, place the unit in full reverse Trendelenburg (foot end down), raise the siderails, and place the I.V. poles and push handles in the up position.
3. Clean the unit with a maximum water temperature of 180 °F (82 °C), maximum air dry temperature (cart washers) is 200 °F (93 °C) for 8 minutes, and maximum water pressure − 1500 psi/103.5 bar.

**Note:** If a handheld wand is being used to wash the unit, the pressure nozzle must be kept a minimum of 24 inches/.61m from the unit. The same water temperatures and stretcher configurations apply as for the cart washer.
CLEANING THE STRETCHER (CONTINUED)

RECOMMENDED CLEANERS

Suggested cleaners for stretchers:
Quaternary Cleaners (active ingredient - ammonium chloride)
Phenolic Cleaners (active ingredient - o-phenylphenol)
Chlorinated Bleach Solution (use 1 part bleach (5.25% sodium hypochlorite) to 100 parts water which equals 520 ppm available chlorine (40 mL of a 5.25% bleach solution per 4000 mL water))

Avoid over-saturation and ensure that the product does not stay wet longer than recommended by the chemical manufacturer’s guidelines for proper disinfecting.

CAUTION

Some cleaning products are corrosive in nature and may cause damage to the product if used improperly. If the products suggested above are used to clean Stryker patient handling equipment, measures must be taken to ensure that the stretcher is wiped with a damp cloth soaked in clean water and thoroughly dried following cleaning. Failure to properly rinse and dry the stretcher will leave a corrosive residue on the surface of the stretcher, possibly causing premature corrosion of critical components. Failure to follow the above directions when using these types of cleaners may void this product’s warranty.

Stretchers must have maintenance performed after a minimum of every fifth washing. See the maintenance manual for specific lubrication instructions.

Do not use abrasive cleaners to clean the display enclosure for the optional scale system. Do not allow cleaning solutions or other fluids to pool on the display unit. Wipe dry all surfaces after spills or cleaning.
CLEANING THE MATTRESS

These instructions are intended to provide recommended cleaning methods for the mattress.

**WARNING**

- Do not immerse mattress in cleaning or disinfectant solutions. Excess moisture could cause equipment malfunction resulting in equipment damage or patient injury.
- Do not allow fluid to pool on the mattress. Fluids can cause corrosion of components and may cause the safety and performance of this equipment to become unpredictable.
- Inspect mattress covers for tears, punctures, excessive wear, and misaligned zippers each time the covers are cleaned. If compromised, the mattress should be removed from service immediately and replaced to prevent cross-contamination.

Stryker mattresses are designed for long-lasting comfort and reliability. The life of the mattress can be adversely affected by an increase in frequency of usage which might include more frequent cleaning and disinfection.

1. Using a clean, soft, damp cloth, wipe down the entire mattress with a mild soap and water solution to remove foreign material.
2. Wipe down the mattress with a clean, dry cloth to remove any excess liquid or cleaning agent.
3. Care must be taken to thoroughly rinse and dry covers following cleaning.
4. Disinfect as necessary with a hospital grade disinfectant after cleaning has been completed. See “Disinfecting the Mattress” on page 69.

**CAUTION**

- Do not iron, dry-clean, or tumble dry the mattress, as this will cause malfunction and damage the product.
- The mattress cover must be completely dry before storing, adding linens or placing a patient on the surface, to prevent the performance of the equipment from being impaired.
- Avoid over exposure to alcohol or hydrogen peroxide. Swelling of the cover material will result.
- Do not allow liquid to seep into the zipper area and watershed cover barrier. Fluids allowed to come in contact with the zipper may leak into the mattress which could impair the equipment performance.

**SPECIAL INSTRUCTIONS**

<table>
<thead>
<tr>
<th>Velcro</th>
<th>To clean and disinfect, saturate with disinfectant, rinse with water, and allow it to evaporate.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soils or Stains</td>
<td>Use neutral soaps and warm water. Do not use harsh cleansers, solvents, or abrasive cleaners.</td>
</tr>
<tr>
<td>Hard-To-Clean Spots</td>
<td>Use standard household/vinyl cleansers and a soft bristle brush on troublesome spots or stains. Pre-soak heavy, dried-on soil.</td>
</tr>
<tr>
<td>Laundering</td>
<td>Laundering is NOT RECOMMENDED. Laundering may substantially decrease the useful life of the mattress.</td>
</tr>
</tbody>
</table>

DO NOT STEAM CLEAN, PRESSURE WASH, HOSE OFF, OR ULTRASONICALLY CLEAN MATTRESSES. Using these methods of cleaning are not recommended and may void this product’s warranty.

**REMOVAL OF IODINE STAINS**

1. Make a solution of 1−2 tablespoons Sodium Thiosulfate in a pint of warm water and use it to wipe the stained area. Clean the stain as soon as possible after it occurs. If stains are not immediately removed, allow solution to soak or stand on the surface before wiping.
2. Rinse surfaces which have been exposed to the solution with clear water before returning the mattress to service.

**Note:** Failure to follow the above directions when using these types of cleaners may void this product’s warranty.
DISINFECTING THE MATTRESS

These instructions are intended to provide recommended disinfecting methods for the mattress.

WARNING

- Disinfect the mattress between patients. Failure to do so could result in cross-contamination and infection.
- Some disinfectants may cause damage to the product if used improperly. If the products described below are used to disinfect the mattress, measures must be taken to ensure the entire surface is wiped with a damp cloth soaked in clean water and thoroughly dried following disinfection. The cover can be damaged when exposed to such disinfectants beyond the manufacturers’ recommendations. Failure to follow these directions when using these types of disinfectants may void this product warranty.
- The mattress cover must be completely dry before storage or adding linens. Failure to remove excess disinfectant could cause degradation of the cover material.

Suggested Disinfectants:

- Quaternaries
- Phenolic Disinfectant
- Chlorinated Bleach Solution (5.25% bleach diluted 1 part bleach to 100 parts water)
- 70% Isopropyl Alcohol

CAUTION

- Frequent or prolonged exposure to higher concentration disinfectant solutions may prematurely age the cover fabric.
- The use of accelerated hydrogen peroxides or quaternaries containing glycol ethers may damage the cover.

1. Ensure surface has been thoroughly cleaned and dried prior to applying disinfectants.
2. Wipe down the mattress with a clean, dry cloth to remove any excess liquid or disinfectant.
3. Care must be taken to thoroughly rinse and dry covers following disinfection.
At a minimum, preventative maintenance should be performed annually. A preventative maintenance program should be established for all Stryker Medical equipment. Preventative maintenance may need to be performed more frequently based on the usage level of the product.

CHECKLIST

_____ All fasteners secure
_____ Siderails move and latch properly
_____ All casters lock with brake pedal engaged
_____ All casters secure and swivel properly
_____ Inspect each caster and remove any wax or debris which may have collected on the caster or braking mechanism
_____ Steer function working properly
_____ Check skins for cracks
_____ Fowler operates and latches properly
_____ Gatch operating properly (Optional equipment)
_____ Trendelenburg/reverse Trendelenburg operates properly from all locations
_____ Ground chain intact
_____ No leaks at hydraulic connections
_____ Hydraulic jacks holding properly
_____ Lubricate where required
_____ Body restraints work properly (Optional equipment)
_____ I.V. pole intact and operates properly (Optional equipment)
_____ Oxygen bottle holder intact and operates properly (Optional equipment)
_____ No rips or cracks in mattress cover
_____ Accessories and mounting hardware in good condition and working properly
_____ Confirm battery powered functionality (Optional equipment)
_____ No cables worn or pinched (Optional equipment)
_____ Power cord and plug are free of damage (Optional equipment)
_____ All electrical connections tight (Optional equipment)
_____ All grounds secure to the frame (Optional equipment)
_____ Ground impedance not more than 200 mΩ (milliohms) (Optional equipment)
_____ Current leakage not more than 300 µA (microamps) (per UL 60601-1) (Optional equipment)
_____ Batteries sufficiently charged (Optional scale system)
_____ Display housing intact and not damaged (Optional scale system)
_____ Load cells intact and not damaged (Optional scale system)
CHECKLIST (CONTINUED)

_____ Foot support knee knob mechanism functions properly and can be secured in place (Optional equipment)
_____ Foot support leg knob mechanism functions properly and can be secured in place (Optional equipment)
_____ Foot support extends to the full position and stops in the correct position (Optional equipment)
_____ Foot support self-tapping screws (6) are functioning properly and are not stripped (Optional equipment)
_____ Scale calibrated properly. Recalibrate, if necessary (Optional scale system)
_____ Upright X-Ray Cassette Holder is working properly
_____ No damage to the Fowler skin and foot skin (Prime X Option)
_____ No damage to the head and foot trays (Prime X Option)

<table>
<thead>
<tr>
<th>Product Serial Number:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<tr>
<td></td>
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<tr>
<td></td>
</tr>
</tbody>
</table>

Completed by: ____________________________________________  Date: _____________________
# Guidance and manufacturer’s declaration - electromagnetic immunity

The Optional Electric Lift/Litter and Optional Scale System is intended for use in the electromagnetic environment specified below. The customer or the user of the Optional Electric Lift/Litter and Optional Scale System should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge (ESD) IEC 61000-4-2</td>
<td>±6 kV contact +8 kV air</td>
<td>±6 kV contact +8 kV air</td>
<td>Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>Electrostatic fast transient/burst IEC 61000-4-4 *</td>
<td>+2 kV for power supply lines +1 kV for input/output lines</td>
<td>+2 kV for power supply lines +1 kV for input/output lines</td>
<td>Main power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5 *</td>
<td>+8 kV differential mode +2 kV common mode</td>
<td>+8 kV differential mode +2 kV common mode</td>
<td>Main power quality should be that of typical commercial and/or hospital environment.</td>
</tr>
<tr>
<td>Voltage dips, voltage variations and short interruptions on power supply input lines IEC 61000-4-11 *</td>
<td>&lt;5%Ut (&gt;95% dip in Ut) for 0.5 cycle 40%Ut (60% dip in Ut) for 5 cycles 70%Ut (30% dip in Ut) for 25 cycles. &lt;5% Ut (&gt;95% dip in Ut) for 5 sec.</td>
<td>&lt;5%Ut (&gt;95% dip in Ut) for 0.5 cycle 40%Ut (60% dip in Ut) for 5 cycles 70%Ut (30% dip in Ut) for 25 cycles. &lt;5% Ut (&gt;95% dip in Ut) for 5 sec.</td>
<td>Main power quality should be that of a typical commercial and/or hospital environment. If the user of the Optional Electric Lift/Litter requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

* Applies to Optional Electric Lift/Litter only

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Note: $U_t$ is the a.c. mains voltage prior to application of the test level.
### Recommended separation distances between portable and mobile RF communications equipment and the Optional Electric Lift/Litter and Optional Scale System.

The Optional Electric Lift/Litter and Optional Scale System is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Optional Electric Lift/Litter and Optional Scale System can help prevent electromagnetic interferences by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Optional Electric Lift/Litter and Optional Scale System as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
</tr>
<tr>
<td></td>
<td>$d=1,2\sqrt{P}$</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance $d$ in meters (m) can be determined using the equation applicable to the frequency of the transmitter, where $P$ is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

**Note 1**

At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

**Note 2**

These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
The Optional Electric Lift/Litter and Optional Scale System is suited for use in the electromagnetic environment specified below. The customer or the user of the Optional Electric Lift/Litter and Optional Scale System should ensure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF IEC 61000-4-6 *</td>
<td>3 Vrms 150 kHz to 80 MHz</td>
<td>3 Vrms</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Optional Electric Lift/Litter and Optional Scale System, including cables, than the recommended separation distance calculated from the equation applicable for the frequency of the transmitter.</td>
</tr>
<tr>
<td>Radiated RF IEC 61000-4-3</td>
<td>3 V/m 80 MHz to 2.5 GHz</td>
<td>3 V/m</td>
<td>Recommended Separation Distance</td>
</tr>
</tbody>
</table>

\[ d = 1.2 \sqrt{P} \]

- 80 MHz to 800 MHz
- 800 MHz to 2.5 GHz

where \( P \) is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and \( d \) is the recommended separation distance in metres (m).

Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.

Interference may occur in the vicinity of equipment marked with the following symbol:

---

**Note 1**
At 80 MHz and 800 MHz, the higher frequency range applies.

**Note 2**
These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

*Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast, and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Optional Electric Lift/Litter and Optional Scale System is used exceeds the applicable RF compliance level above, the Optional Electric Lift/Litter and Optional Scale System should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Optional Electric Lift/Litter and Optional Scale System.

**Note 2**
Over the frequency range 150 kHz to 80 MHz, field strengths are less than 3 V/m.

* Applies to Optional Electric Lift/Litter only
**Guidance and manufacturer’s declaration - electromagnetic emissions**

The Optional Electric Lift/Litter and Optional Scale System is intended for use in an electromagnetic environment specified below. The customer or the user of the Optional Electric Lift/Litter and Optional Scale System should ensure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions CISPR 11</td>
<td>Group 1</td>
<td>The Optional Electric Lift/Litter and Optional Scale System uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF Emissions CISPR 11</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>Harmonic Emissions IEC 61000-3-2</td>
<td>Class A</td>
<td>The Optional Electric Lift/Litter and Optional Scale System is suitable for use in all establishments other than domestic and those directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Voltage Fluctuations Flicker Emissions IEC 61000-3-3</td>
<td>Complies</td>
<td></td>
</tr>
</tbody>
</table>

* Applies to Optional Electric Lift/Litter only
LIMITED WARRANTY

Stryker Medical Division, a division of Stryker Corporation, warrants to the original purchaser the Stryker Model 1105 Prime Series stretcher to be free from defects in material and workmanship for a period of two (2) years after date of delivery. Stryker's obligation under this warranty is expressly limited to supplying replacement parts and labor for, or replacing, at its option, any product which is, in the sole discretion of Stryker, found to be defective. If requested by Stryker, products or parts for which a warranty claim is made shall be returned prepaid to the factory. Any improper use or any alteration or repair by others in such manner as in Stryker's judgment affects the product materially and adversely shall void this warranty. Any repair of Stryker products using parts not provided or authorized by Stryker shall void this warranty. No employee or representative of Stryker is authorized to change this warranty in any way.

Stryker Medical Stretcher products are designed for a 10 year expected service life under normal use, conditions, and with appropriate periodic maintenance as described in the maintenance manual for each device. Stryker warrants to the original purchaser that the welds on its Stretcher products will be free from structural defects for the expected 10 year life of the Stretcher product as long as the original purchaser owns the product.

This statement constitutes Stryker’s entire warranty with respect to the aforesaid equipment. Stryker makes no other warranty or representation, either expressed or implied, except as set forth herein. There is no warranty of merchantability and there are no warranties of fitness for any particular purpose. In no event shall Stryker be liable here under for incidental or consequential damages arising from or in any manner related to sales or use of any such equipment.

Warranty does not include any disposable items, I.V. poles (except for Stryker permanently attached poles), mattresses, batteries, or damage resulting from abuse.

TO OBTAIN PARTS AND SERVICE

Stryker products are supported by a nationwide network of dedicated Stryker Field Service Representatives. These representatives are factory trained, available locally, and carry a substantial spare parts inventory to minimize repair time. Simply call your local representative, or call Stryker Customer Service USA at 1-800-327-0770.

SERVICE CONTRACT COVERAGE

Stryker has developed a comprehensive program of service contract options designed to keep your equipment operating at peak performance at the same time it eliminates unexpected costs. We recommend that these programs be activated before the expiration of the new product warranty to eliminate the potential of additional equipment upgrade charges.

A Service Contract helps to:

- Ensure equipment reliability
- Stabilize maintenance budgets
- Diminish downtime
- Establish documentation for JCAHO
- Increase product life
- Enhance trade-in value
- Address risk management and safety
Warranty

SERVICE CONTRACT PROGRAMS

Stryker offers the following service contract programs:

<table>
<thead>
<tr>
<th>Service Agreement Options</th>
<th>Premium</th>
<th>Complete</th>
<th>Standard *</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annually scheduled preventative maintenance</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All parts**, labor, and travel</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Unlimited emergency service calls</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Priority one contact: two hour phone response</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Most repairs will be completed within 3 business days</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>JCAHO documentation</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>On-site record of PM &amp; emergency service</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Factory-trained Stryker service technician</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Stryker authorized parts used</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Service during regular business hours (8-5)</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

* Replacement parts and labor for products under PM contract will be discounted.
** Does not include any disposable items, I.V. poles (except for Stryker permanently attached poles), mattresses, batteries, or damage resulting from abuse.

Stryker Medical also offers personalized service contracts.

Pricing is determined by age, location, model and condition of product.

For more information on our service contracts, please call your local representative.

RETURN AUTHORIZATION

Merchandise cannot be returned without approval from the Stryker Customer Service Department. An authorization number will be provided which must be printed on the returned merchandise. Stryker reserves the right to charge shipping and restocking fees on returned items. Special, modified, or discontinued, items not subject to return.

DAMAGED MERCHANDISE

ICC Regulations require that claims for damaged merchandise must be made with the carrier within fifteen (15) days of receipt of merchandise. Do not accept damaged shipments unless such damage is noted on the delivery receipt at the time of receipt. Upon prompt notification, Stryker will file a freight claim with the appropriate carrier for damages incurred. Claim will be limited in amount to the actual replacement cost. In the event that this information is not received by Stryker within the fifteen (15) day period following the delivery of the merchandise, or the damage was not noted on the delivery receipt at the time of receipt, the customer will be responsible for payment of the original invoice in full. Claims for any short shipment must be made within thirty (30) days of invoice.

INTERNATIONAL WARRANTY CLAUSE

This warranty reflects U.S. domestic policy. Warranty outside the U.S. may vary by country. Please contact your local Stryker Medical representative for additional information.
Recycling Passport

Assembly part number:
1070-110-260 (Reference Only)
1070-110-265 (Reference Only)
1070-110-360 (Reference Only)
1070-110-365 (Reference Only)
1070-110-270 (Reference Only)
1070-110-370 (Reference Only)

<table>
<thead>
<tr>
<th>Item</th>
<th>Recycling/Material Code</th>
<th>Important Information</th>
<th>Qty</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-037-820) Scale Control Non-Backlit Keypad Assembly</td>
<td></td>
<td>1</td>
</tr>
</tbody>
</table>
Recycling Passport

Assembly part number:
1070-117-600 (Reference Only) - 26"
1070-117-300 (Reference Only) - 30"

<table>
<thead>
<tr>
<th>Item</th>
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<tbody>
<tr>
<td>A</td>
<td>(1008-037-057) Load Cell</td>
<td></td>
<td>2</td>
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1070-110-265 (Reference Only)
### Recycling Passport

Assembly part number: 1070-237-020 (Reference Only)

<table>
<thead>
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<th>Item</th>
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<th>Qty</th>
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<tbody>
<tr>
<td>A</td>
<td>(1008-037-830) Scale Control Assembly</td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>B</td>
<td>(1070-137-029) AA Battery Assembly</td>
<td></td>
<td>1</td>
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</table>
Recycling Passport

Assembly part number: 1008-010-302 (Reference Only)

<table>
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<tr>
<th>Item</th>
<th>Recycling/Material Code</th>
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<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-015-820) Patient</td>
<td>Lockout Assembly</td>
<td>1</td>
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</table>
Assembly part number:
1008-011-320 (Reference Only) - Right
1008-011-330 (Reference Only) - Left

<table>
<thead>
<tr>
<th>Item</th>
<th>Recycling/Material Code</th>
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<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-011-016) Siderail Keypad, Right (1008-011-017) Siderail Keypad, Left</td>
<td></td>
<td>1</td>
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</table>
Recycling Passport

Assembly part number: 1008-015-020 (Reference Only)

<table>
<thead>
<tr>
<th>Item</th>
<th>Recycling/Material Code</th>
<th>Important Information</th>
<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-015-800) Staff Control Assembly</td>
<td></td>
<td>1</td>
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</tbody>
</table>
Assembly part number: 1070-010-100 (Reference Only)

<table>
<thead>
<tr>
<th>Item</th>
<th>Recycling/Material Code</th>
<th>Important Information</th>
<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(0058-135-000) Li-ION Smart Battery Pack</td>
<td></td>
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</tr>
<tr>
<td>B</td>
<td>(1008-237-850) Scale Control Assembly</td>
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</table>
Assembly part number: 1070-010-100 (Reference Only)

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>A</td>
<td>(1008-037-810) Scale Control Backlit Keypad Assembly</td>
<td></td>
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</table>
### Recycling Passport

Assembly part number: 1070-010-200 (Reference Only)

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<th>Item</th>
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<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(0058-134-000) Li-ION Smart Battery Pack</td>
<td></td>
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<tr>
<td>B</td>
<td>(1008-237-840) Scale/Chaperone Control Assembly</td>
<td></td>
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</table>
Assembly part number: 1070-010-200 (Reference Only)

<table>
<thead>
<tr>
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<th>Qty</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-037-800) Scale/Chaperone Keypad Assembly</td>
<td></td>
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</table>
Assembly part number: 1008-005-500 (Reference Only)

<table>
<thead>
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<th>Qty</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-002-800) Non-Lift Control PCB Assembly</td>
<td></td>
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</table>
Assembly part number: 1008-005-510 (Reference Only)

<table>
<thead>
<tr>
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<th>Qty</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(1008-002-810) Lift Control PCB Assembly</td>
<td></td>
<td>1</td>
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</table>